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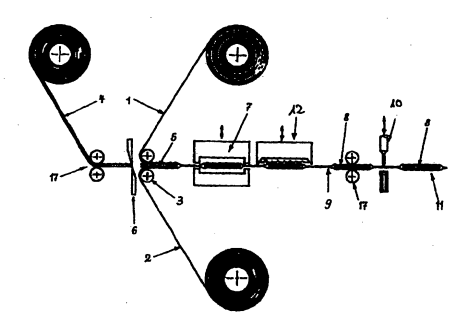
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- (51) Int.Cl.⁶ B65D 75/42
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- (54) UNITE D'EMBALLAGE PRIMAIRE POUR FORMES D'ADMINISTRATION DU TYPE FILM OU PLAQUETTE
- (54) PRIMARY PACKAGING UNIT FOR FILM-LIKE OR OBLATE-LIKE ADMINISTERED SHAPES



- (57) L'invention concerne une unité d'emballage primaire pour formes d'administration du type film ou plaquette, destinées à l'administration par voie orale et comportant chacune une section constituée d'une bande supérieure (1) et d'une bande inférieure (2) de matière d'emballage. L'invention se caractérise en ce
- (57) The invention relates to a primary packaging unit for film-like or oblate-like administered shapes used for oral application. The packaging unit each comprises a section of a packing upper (1) and lower (2) web. The invention is characterized in that a plurality of dosing units (5) of a film-like or oblate-like administered

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que plusieurs doses unitaires (5) d'une forme d'administration du type film ou plaquette sont scellées, à une certaine distance les unes des autres, individuellement, dans une unité d'emballage primaire, dans des compartiments (8) plats réalisés sans formage à froid ou à chaud du matériau d'emballage, et en ce que des perforations se trouvent entre lesdits compartiments (8), lesquelles permettent la

séparation de compartiments (8) individuels en fonction

shape are arranged at a distance from one another and are individual sealed inside compartments (8) of a primary packaging unit. Said compartments are produced without cold or heat forming the packing material. The invention is also characterized by having perforations which are located between compartments (8), said perforations permitting the separation of individual compartments (8) according to need.

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Veröffentlicht

Mit internationalem Recherchenbericht. Vor Ablauf der für Änderungen der Ansprüche zugelassenen Frist; Veröffentlichung wird wiederholt falls Änderungen eintreffen.

- (54) Title: PRIMARY PACKAGING UNIT FOR FILM-LIKE OR OBLATE-LIKE ADMINISTERED SHAPES
- (54) Bezeichnung: PRIMÄRVERPACKUNGSEINHEIT FÜR FILM- ODER OBLATENARTIGE DARREICHUNGSFORMEN

(57) Abstract

The invention relates to a primary packaging unit for film-like or oblate-like administered shapes used for oral application. The packaging unit each comprises a section of a packing upper (1) and lower (2) web. The invention is characterized in that a plurality of dosing units (5) of a film-like or oblate-like administered shape are arranged at a distance from one another and are individual sealed inside compartments (8) of a primary packaging unit. Said compartments are produced without cold or heat forming the packing material. The invention is also characterized by having perforations which are located between compartments (8), said perforations permitting the separation of individual compartments (8) according to need.

(57) Zusammenfassung

Eine Primärverpackungseinheit für film- oder oblatenartige Darre-

ichungsformen zur oralen Applikation mit je einem Abschnitt einer Packstoffober- (1) und -unterbahn (2) ist dadurch gekennzeichnet, dass mehrere Dosiseinheiten (5) einer film- oder oblatenartigen Darreichungsform im Abstand zueinander einzeln in flachen, ohne Kaltoder Heissverformung des Packmaterials erzeugten Kompartimenten (8) eingesiegelt in einer Primärverpackungseinheit vorliegen, und sich zwischen Kompartimenten (8) Perforationen befinden, welche die bedarfsweise Abtrennung einzelner Kompartimente (8) ermöglichen.

The present invention relates to a primary packaging unit for film-like or wafer-like administration forms for oral application. The invention especially relates to a primary packaging unit which is formed out of the film-like or wafer-like administration form to be packaged as well as a section of an upper and a lower web of packaging material, respectively.

Film-like or wafer-like administration forms for oral application are known e.g. from the patents or applications US 3 007 848, DE 24 32 925, DE 27 46 414 and EP 219 762. Said administration forms differ from conventional solid application forms such as tablets or capsules especially in their geometrical form and their production. They all have a thin, flat-shaped form, whereby differences with regard to flexibility, brittleness, smoothness or consistency can lead to either film- or sheet-like, paper-like, or wafer-like characteristics. For the production of said administration forms, the extrusion and coating processes applied in industrial film production were especially recommended.

Depending on the purpose of application, two basic types of embodiment suggest themselves. The first type comprises variations with rapid disintegration or rapid release for disintegration in the oral cavity immediately upon application under release of an active substance, whereby the term "rapid disintegration" in the sense of this invention relates to a disintegration time of several seconds up to a maximum of several minutes under influence of saliva. The other type comprises variations which disintegrate slowly or practically not at all and are especially suited for slow and continuous active substance

release and which, through addition of mucoadhesive materials, are able to adhere to the oral mucosa during the release of active substance. Both of these basic types can be embodied so that, depending on the incorporated active substance, they are suited for a local therapy of the oral mucosa or the systematic application of active substances.

The packaging of these administration forms in primary packaging units cannot simply be carried out with the usual processes, packaging means or machines commonly used for conventional pharmaceutical products such as capsules or tablets. A primary packaging unit for solid administration forms in individual doses, embodied from a modern point of view, should on the one hand protect the product from outer influences and on the other hand enable the deliberate and controllable removal of a single dosage unit at the desired time of intake, whereby the removal of the dosage unit from the primary packaging unit should be carried out in such a way that the administration form is not damaged.

Whereas tablets and capsules are often filled into glasses or boxes in larger amounts, which certainly would not suffice to fulfil the above requirements, it is in many cases customary to package dosage units in blister packages or deep-drawn packages. Such primary packaging units contain a plurality of dosage units which are each individually sealed in a cavity between two sections of packaging material webs. The cavity is created through hot or cold forming of the lower web of packaging material with the help of an adequate tool before filling. After the cavities are filled, the upper web of packaging material is supplied and sealed together with the lower web of packaging material.

In modern blister packages, the dosage units are removed by exerting a pressure, with a finger, on the outer side of the deformed areas of the lower packaging material web and

thus on the tablet or capsule contained in the cavity created through deformation, whereby the exerted pressure is sufficient to break through the upper web of packaging material and press out the dosage unit. This is, however, only possible if the material of the upper packaging material web does not exceed a certain strength.

This concept for primary packaging units has become widely known and used for conventional administration forms. For administration forms with film-like or wafer-like embodiments, however, it presents considerable disadvantages. In experiments carried out to this effect, two disadvantages have proved to be particularly serious, one of which concerns the production and the other of which concerns the removal of dosage units from primary packaging units of this kind.

Film-like or wafer-like oral administration forms especially the rapid release kinds - are generally very much lighter and less compact than conventional tablets or capsules. The recommended dimensions of film-like or waferlike dosage units are approximately from 1 cm2 (e.g. DE 27 46 414) to 3 cm² or more (e.g. DE 24 32 925), with a thickness of approximately 0.05 to 1 mm (e.g. DE 24 32 925). Using common pharmaceutical base materials, this then results in dosage units with a mass of approximately 5 to 100 mg, whereby the typical and preferred embodiments would tend to lie in the lower margin of this span. It has turned out, however, that enclosing such thin films or wafers in blisters is quite problematic. Especially in the case of higher machine speeds, air movement caused by moving machine parts and often also electrostatic charging of the packaging materials lead to the result that the dosage units cannot be correctly positioned in the blister or are wafted back out of the blister after being positioned therein. Although it is quite possible to produce deepdrawn primary packaging units with oral films or wafers, this is a complicated and inefficient packaging concept due to the problems mentioned above. The removal of film-like or wafer-like administration forms from blister packages which correspond to the conventional primary packaging for tablets and capsules is also problematic. A flat-shaped dosage unit lying in a cavity can hardly be pressed through the material of the upper web of packaging material; it has neither the necessary format nor the mechanical strength. The danger of damaging the dosage unit while pressing it out of the package is relatively large. Even if one first tries to break the material of the upper packaging material web in another way, e.g. using a fingernail, it is not easy to grip and remove a flat dosage unit in the exposed cavity, except when very large cavities are chosen, which is disadvantageous because of other reasons such as the enclosed air space, which is too large in relation the small mass of the administration form.

The use of conventional packaging means results in additional difficulties if the film-like or wafer-like administration forms are rather brittle and fragile. In this case, a dimensionally stable blister package can offer a certain amount of product protection during storage, but it makes the removal of the dosage units even more difficult.

In addition to these disadvantages of conventional blister packages for film-like or wafer-like administration form, the choice of adequate packaging materials for blister packages is limited; also, the available materials do not belong to the especially cheap packaging materials.

Several approaches for the creation of a primary packaging unit for film-like or wafer-like administration forms without the above described disadvantages of the state of the art are found in US 3 007 848. The solutions presented

in this document are partially of interest for all filmlike and wafer-like administration forms although US 3 007 848, in contrast to the present invention, in the narrower sense refers to (1) wafers produced through extrusion or through printing of edible films, whereby (2) said wafers are not intended for application in the oral cavity, but rather for swallowing, and (3) are for this purpose optionally first sealed into film strips of an edible, smooth and easily swallowable film. The cited document does, however, teach the packaging of wafers by sealing the dosage units between two films which can in a general sense be understood as packaging materials. In addition, it teaches the only "light" sealing of the dosage units to enable an easier opening of the compartments and removal of the wafers. Finally, it also teaches an unsealed outer area of the packaging material which facilitates the gripping of the packaging material films and their pulling apart to remove the wafers.

US 3 007 848, which constitutes a state-of-the-art solution near to the present invention, does not, however, fulfil all requirements for an adequate primary packaging unit for film-like or wafer-like administration forms; several disadvantages and problems remain unsolved or newly arise through the embodiment of the packaging unit suggested therein.

On the one hand, the packaging units suggested therein contain only one wafer each — disregarding the intermediary product, which comprises an undefined but very large amount of packaged dosage units as a sort of tape goods that can be rolled up. A practicable primary packaging unit should, however, for various reasons generally contain a clearly defined amount of dosage units. If this requirement is not fulfilled, clear disadvantages arise for the secondary packaging: first, the small primary packaging units

separated according to US 3 007 848 must be filled with one wafer each, collected, counted to package sizes of e.g. 20 units and gathered together, which costs a considerable effort and leads to unwieldy secondary packaging formats. For each later extraction, a primary packaging unit would have to be removed, opened, and the wafer extracted, whereby a control of the intake up to a certain point in time is very difficult. In the case of a secondary packaging unit with 50 wafers, e.g., it will hardly be possible, without an arduous counting of the remaining wafers, to keep track of whether a certain due dose has already been taken or not.

It is thus the object of the present invention to provide a primary packaging unit for film-like or wafer-like administration forms which fulfils all of the above mentioned requirements without having the above described disadvantages of the state of the art.

This object is achieved by providing a primary packaging unit for film-like or wafer-like administration forms for oral application with a section of an upper and a lower web of packaging material, respectively. Said packaging unit is characterized in that a plurality of dosage units of a film-like or wafer-like administration form, individually sealed in flat compartments formed without cold or hot forming of the packaging material and spaced at a distance to one another, are present in a primary packaging unit, and in that there are perforations between the compartment which enable the separation of individual compartments, if necessary.

This combination of characteristics is necessary to achieve the claimed, practicable primary packaging unit. A simple variation of the concept of US 3 007 848 to the effect that the intermediate product, which is present e.g. as rolled stock or tape goods, is cut - not after each wafer, as

claimed, but rather after every e.g. tenth wafer - does not suffice to achieve the object stated above. A thus produced packaging unit would contain a defined amount of dosage units; these could, however, not be extracted easily and without problems. Experiments have shown that when opening such a package for the extraction of one dosage unit, the sealed seams or sealed areas around several dosage units adjacent to this unit are generally opened simultaneously, so that several dosage units are exposed and no longer protected by the primary packaging. As described above, the targeted removal of a single dosage unit by pressing it through the primary packaging unit is not possible either because of the low mechanical strength of the administration form in relation to the primary packaging material.

It was found that a primary packaging unit which satisfactorily fulfils the objects of the invention must also have an additional characteristic: a perforation between the compartments in which the individual dosage units are situated, whereby said perforation must be such that it is possible, for the extraction of a single dosage unit, to first separate the compartment containing this dosage unit from the primary packaging unit, if necessary, so that the compartment can then be opened without damaging further compartments. The perforation further offers the advantage that in a respective embodiment with ideally only a few small holding points, it also enables the targeted opening of a compartment without first detaching it from the primary packaging unit, without simultaneously opening further compartments.

A further advantage of the primary packaging unit according to the invention is the relatively small head space of the compartments in which the dosage units are situated. Oxidation- or moisture-sensitive products can thus largely be protected against the harmful influences of atmospheric oxygen and air humidity if the primary packaging materials are chosen accordingly.

A further advantage of the primary packaging unit according to the invention is the small amount of required packaging material and the compact, space-saving format. A folding box with a height of 1 cm, e.g., can easily hold ten or more primary packaging units with ten dosage units each.

A further advantage of the primary packaging unit according to the invention is the possibility of using, for the lower web of packaging material, materials which are considerably thinner and cheaper than those which are suitable for the production of blister packages and for cold and hot forming and which must have a certain minimum thickness and thus also a minimum weight.

A further advantage of the primary packaging unit according to the invention is the possibility of visually presenting therapy patterns on the package by means of printing. Thus, a packaging unit can e.g. be embodied as a 7-day-package with seven dosage units of a drug which is to be taken once a day, whereby the individual compartments of the packaging unit are printed with the names or abbreviations of the different days of the week. With the help of this therapy pattern printed onto the package, patients can very easily control their intakes. In a preferred embodiment, the subject matter of the invention contains printing.

Because film-like or wafer-like administration forms, as described e.g. in DE 24 32 925, are especially advantageously first produced as a cast film from which the dosage units can be obtained by cutting or punching, a further preferred embodiment of the primary packaging unit according to the invention contains dosage units which are sections or punched-out pieces of cast films. Cast films in

the sense of this invention include all film-like compositions produced by casting of carrier materials or coating of the same with polymer-containing solutions, suspensions or emulsions, and subsequent drying.

A further preferred embodiment of the primary packaging unit according to the invention contains sealed seams or sealed areas between the sections of the upper and lower webs of packaging materials which are peelable. The term peelable in the sense of this invention comprises all sealed seams and sealed areas which can be separated with a moderate pulling force, e.g. less than approx. 10 N/15 mm, whereby the packaging material web sections generally remain intact. For the production of such peelable sealing seams, special sealing materials, e.g. so-called "peel-PE" a special polyethylene that generally contains a further polymer such as e.g. polystyrene, and special sealing conditions (pressure, time, temperature) are used. It is, however, also possible to seal conventional sealing materials under such conditions that the result is not a composite in the form of a melted sealed seam but rather a peelable seam.

A further preferred embodiment of the primary packaging unit according to the invention provides that next to each compartment, outside of the sealed areas or sealed seams, there is an unsealed edge on at least one side. This unsealed edge serves as a gripping tab for an easy gripping of the sections of the upper and lower webs of packaging material and separation of the packaging materials to open a compartment. In a further preferred embodiment, these gripping tabs or unsealed edges have different respective lengths for the sections of the upper and lower webs of packaging material. If one of these two packaging material web sections protrudes at the edge, it is especially easy to grip and bend away from of the second packaging material

web section, due to which this second section can also be gripped more easily.

Packaging material webs for the production of primary packaging units according to the invention can be single-layered; generally, however, they will be multi-layered to be able to meet the requirements that must be posed to modern packaging materials and in connection with film-like or wafer-like administration forms.

Common often-used layers are e.g. kraft paper for providing rigidity, plastic films for providing tensile strength and tightness of the packaging material, sealing lacquers for a better sealing capacity, protective lacquers for impregnation of the kraft paper, aluminum for an especially high tightness, glue for the cohesion of individual layers, etc.. In terms of economic considerations, optimized packaging material laminates do not have more layers or greater layer thicknesses than necessary for the respective object.

In certain cases, it will be necessary to employ a certain packaging material laminate for both the upper and the lower web of packaging material for a primary packaging unit according to the invention. If, e.g., an especially high impermeability to gas is necessary which can only be achieved by means of an aluminum barrier layer, it will be necessary to use this element in both packaging material webs.

In other cases, however, different requirements can be posed to the upper web and the lower web. If, e.g., a primary package is to have a certain minimum rigidity – for better handling, a preferred embodiment of the primary packaging unit according to the invention employs a web of packaging material with a bending rigidity of at least x in the case of a combined minimum strength of y μ m – it is

sufficient if this rigidity is mainly provided by one of the packaging material webs, while the other web of packaging material can be optimized under consideration of other economic or technical factors.

A further preferred variation of the primary packaging unit according to the invention with two differently embodied webs of packaging material contains a transparent upper packaging material web section through which the dosage units of the administration form can be seen through the intact package. The definition of upper and lower web is arbitrary; if one transparent and one non-transparent web of packaging material is used, the transparent web is herewith defined as upper web in the sense of this invention. One of the advantages of this variation is the easy visual controllability of the compartments or the dosage units and their condition. A further advantage is that through a transparent upper web, printing on the upper surface of the lower web or also on the dosage units can be discerned. As such printing offers advantages e.g. with regard to intake control, as described above, a preferred embodiment of the primary packaging unit according to the invention contains a transparent upper packaging material web section and either a lower web section printed on its upper surface or dosage units printed on their upper side.

Packaging units according to the present invention are suited for all state-of-the-art film-like or wafer-like administration forms. These include simple, single-layered preparations which generally disintegrate rapidly in saliva, as well as multi-layered systems which adhere to the mucosa and release their active substance over a longer amount of time and the layers of which accordingly have different compositions, whereby at least one layer disintegrates only slowly or not at all in saliva and a further layer has mucoadhesive characteristics.

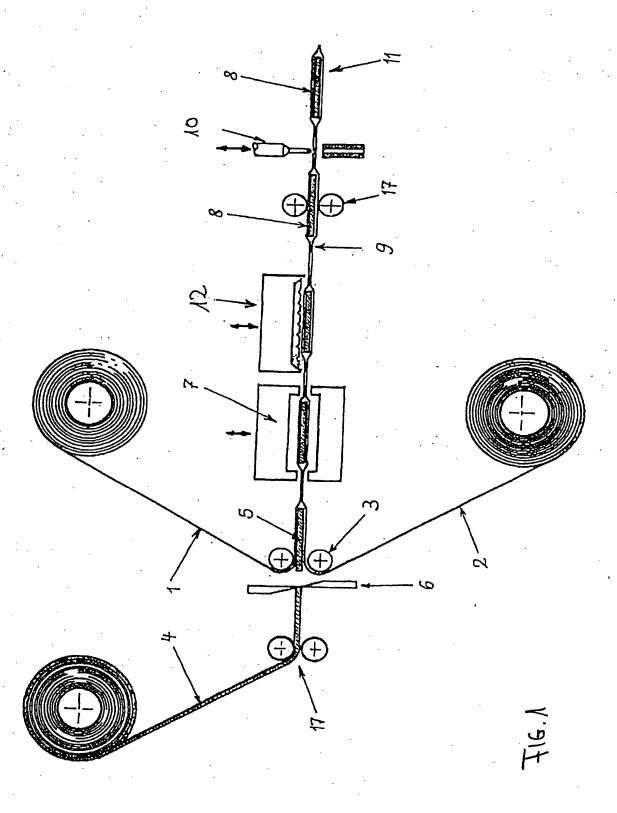
Primary packaging units according to the invention can be machine manufactured with surprising efficiency. A preferred production process, shown schematically in Fig. 1, for packaging units with square or rectangular dosage units (5) comprises at least the following fundamental process steps which can, if necessary, be supplemented by further steps for printing, additional forming of the packaging units, etc.: in a first step, an upper web of packaging material (1) and a lower web of packaging material (2) without cold or hot forming are conveyed on top of one another by means of respective deflecting rolls (3), whereby the film-like or wafer-like administration form (4) is simultaneously conveyed between the two packaging material webs with the help of pulling devices (17) in the form of rolls or tongs. It is advantageous if the film-like or wafer-like administration form is already provided as a web material - single-webbed or multi-webbed, parallel and spaced at a distance to one another - with the desired width of the dosage units (5). It is also advantageous if the diameter of the deflecting rolls is smaller than the length of the dosage units in the conveying direction of the webs. In a further process step, individual dosage units (5) are singled out of the webformed administration form by means of a cross-cutting apparatus (6) which is positioned immediately in front of the deflecting rolls. In a further process step, the two webs of packaging material are sealed together with the help of a heated sealing tool (7) in such a way that the single dosage units (5) are sealed into compartments (8) and are completely enclosed by sealed seams or sealed areas In a further process step, perforations are punched between the compartments (8) by means of a punching device (12). In a further process step, primary packaging units (11) can be partitioned off by means of a second crosscutting or punching device.

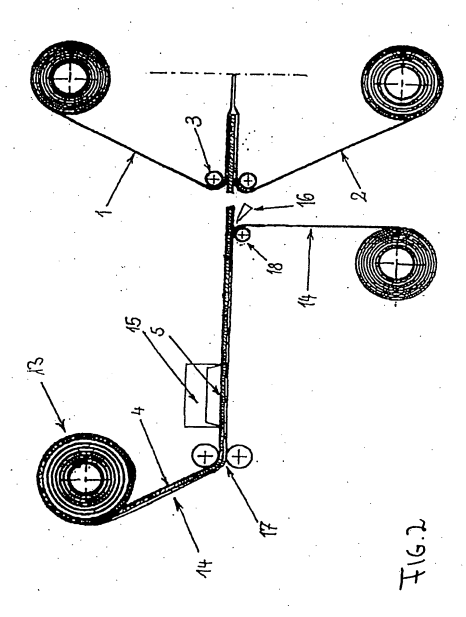
Especially if dosage units (5) are desired which do not have a rectangular or square geometrical form, another multi-step production process is preferred which is schematically shown in Fig. 2. The process steps described here can also be supplemented by further steps or varied in their order if necessary. In one process step, the process comprises providing a laminate (13) of the web-formed, film-like or wafer-like administration form (4) and a carrier sheet (14), out of which the dosage units (5) are punched with a punching device (15) in a further process step, whereby the carrier sheet (14) is not punched through. In a further process step, the punched laminate (13) is rerouted over an edge or a deflecting roll (18) with the help of pulling devices in the form of rolls or tongs (17) so that the dosage units (5) thereby become detached from the carrier sheet (14). If necessary, an additional stripping device (16) can be used for this. In a further process step, an upper web of packaging material (1) and a lower web of packaging material (2) without cold or hot forming are conveyed on top of one another by means of respective deflecting rolls (3), whereby the dosage units (5) becoming detached from the carrier sheet (14) are simultaneously conveyed between the two webs of packaging material (1, 2). In a further process step, the two webs of packaging material are sealed together with the help of a heated sealing tool (7) according to Fig. 1 in such a way that the single dosage units (5) are sealed into compartments (8) and are completely enclosed by sealed seams or sealed areas (9). In a further process step, perforations are punched between the compartments (8) by means of a punching device (12). In a further process step, primary packaging units (11) can be partitioned off by means of a second cross-cutting or punching device.

Claims

- process for manufacturing a primary packaging unit for film-like or wafer-like administration forms for oral application with a section of an upper web of packaging material (1) and of a lower web of packaging material (2), with a plurality of dosage units (5) being sealed in flat compartments (8), and perforations being provided between said compartments, characterized in that the film-like or wafer-like administration form is provided by a web-like laminate comprising a carrier sheet, that the dosage units are punched out of the laminate and subsequently the punched-out laminate (13) is advanced and diverted in such a way that the dosage units (5) become detached from the carrier sheet (14) and are led between the packaging material webs (1, 2), and that subsequently the latter (1, 2) are sealed to each other in sections in such a way that compartments (8) comprising dosage units (5) are formed.
- 2. Process according to Claim 1, characterized in that for partitioning off of the dosage units, the laminate (13) is drawn from a supply roll by means of pulling devices (17) in the form of rolls or tongs, is punched and led around the edge or the deflecting roll (18).
- 3. Process according to Claim 1, characterized in that the webs of packaging material (1, 2) are conveyed on top of one another by means of one deflecting roll (3) per web while simultaneously the dosage units (5) which are becoming detached from the carrier film (14) are pushed between the two webs of packaging material (1, 2).

4. Process according to Claim 1, characterized in that in a further step, individual primary packaging units (11) are severed by means of a cross-cutting or punching device.





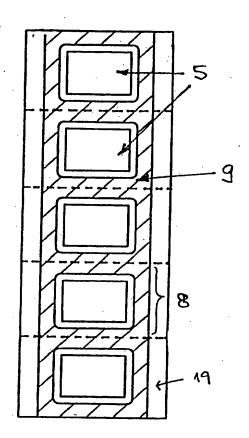
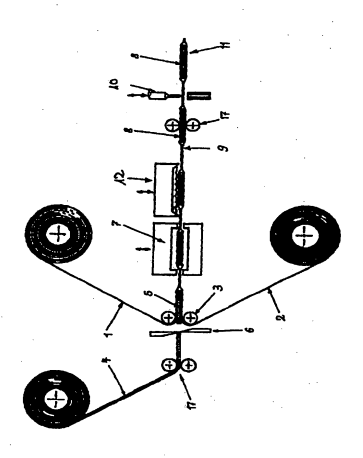


FIG.3



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